Exhibit 4

(22) Application Date: 2003.09.26

(19) AUSTRALIAN PATENT OFFICE

(54) Title Recombinant super-compound interferon

 $(51)^6$ International Patent Classification(s)

A61K 038/21 A61P 031/12 C12N 015/63 A61P 001/16 C12N 015/20 C12N 015/70

Application No: 2003248419 (43)Publication Date : 2003.11.06

(43)Publication Journal Date: 2003,11.06

Divisional of: (62)

2002235707

(21)

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Patents Act 1990

COMPLETE SPECIFICATION
FOR A STANDARD PATENT
ORIGINAL

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Title: RECOMBINANT SUPER-COMPOUND INTERFERON

Associated Provisional Applications: No(s).:

The following statement is a full description of this invention, including the best method of performing it known to me/us:

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COMS ID No: SMBI-00433101 Received by IP Australia: Time (H:m) 18:11 Date (Y-M-d) 2003-09-26

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RECOMBINANT SUPER-COMPOUND INTERPERON

The application is a continuation-in-part application of International Patent application No. PCT/CNOZ/00128, filed on 28 February 2002, which claims priority of Chinese Application No. 01104357.9, filed on 28 February 2001, the contents of which are incorporated by reference here into this application.

Throughout this application, warious references are referred to. Disclosures of these publications in their entireties are hereby incorporated by reference into this application in order to more fully describe the state of the art to which this invention pertains.

FIELD OF THE INVENTION

This invention is related to a recombinant super-compound interferon (rSIN-co) with changed spatial configuration. One characteristic of TSIN-co in this invention is that it cannot only inhibit DNA (deoxyribonucleic acid) duplication of the Hepatitis B virus but also the secretion of HBsAg and HBsAg.

25 BACKGROUND OF THE INVENTION

rSIFN-co is a new interferon molecule constructed with the most popular conservative entine acid found in natural human a-IFN subtypes using genetic engineering methods. Onited States Patent Nos. 4,695,623 and 4,897,471 have described it. rSIFN-co had been proved to have broad-spectrum IFN activity and virus- and tumor-inhibition and natural Killer cell activity. United States Patent No. 5,372,808 by Amgen. Inc. addresses treatment rSIFN-co. Chinese Patent No. 97193506,8 by Amgen, Inc. addresses re-treatment of rSIFN-co on Hepatitis C. Chinese Patent No. 98114663.5 by Shenzhen Jusheng Bio-engineering Ltd. addresses treatment

of rSIFN-co on Hepatitis B and Hepatitis C.

The United States Food and Drug Administration (FDA) authorized Ammgen to produce rSIFN-co with E. Coli. for clinical Repetitis C treatment at the end of 1997.

Hepatitis B patients can be identified when detecting HBsAg and the HBsAg. a-TPN is commonly used in clinics to treat Hepatitis B. IFN binds superficial cell membrane raceptors, inhibiting DNA and RNA (ribonucleic acid) duplication, including inducing some enzymes to prevent duplication of the virus in hepatitie-infected cells. All FPNs can inhibit only the DNA duplication of viruses, not the e and s antigen.

This disclosure describes recombinant super-compound interferon, method to produce the same and uses thereof.

The above references to and descriptions of prior proposals or products are not intended to be, and are not to be construed as, statements or admissions of common general knowledge in the art in Australia.

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SUMMARY OF THE INVENTION

This invention provides a recombinant super-compound interferon or an equivalent thereof with changed spatial An equivalent is a molecule which is similar in function to the super-compound interferon. The super-compound interferon possesses anti-viral or antitumor activity. This invention also provides an artificial gene codes for the super-compound interferon or its equivalent.

This invention provides a process for production of super-compound interferon comprising recombinant introducing an artificial gene with selected codon preference into an appropriate host, culturing said introduced host in an appropriate condition permitting expression of said super-compound interferon and harvesting the expressed super-compound interferon.

This invention provides a composition comprising the 20 recombinant super-compound interferon or its equivalent and a suitable carrier. This invention further provides a pharmaceutical composition comprising the recombinant super-compound interferon or its equivalent and a pharmaceutically acceptable carrier. 25

> This invention provides a method for treating viral diseases or tumor in a subject comprising administering to the subject an effective amount of the super-compound interferon or its equivalent.

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This invention provides the above-described method wherein super-compound interferon was administered via oral, vein muscle injection, peritoneal subcutaneous injection, masal, mucosal administration, by inhalation via an inspirator.

DETAILED DESCRIPTION OF THE FIGURES

- 5 Figure 1. rSIFN-co cDNA sequence designed according to E. Colf. codon usage and deduced rSIFN-co amino acid sequence
 - Figure 2. Sequence of another super-compound interferon
- 10 Figure 3. Diagram of plac T7 cloning vector plasmid
 - Figure 4. Diagram of pHY-4 expression vector plasmid
 - Figure 5. Construction process of expression plasmid pHY-5
- 73 Zigure 6-A. Circular Dichroism spectrum of Infergen Spectrum range: 250nm 190nm Sensitivity: 2 m°/cm

Light path: 0.20 cm

- Equipment: Circular Dichroism J-500C

 Samples: contains 30µg/ml IFN-con1, 5.9 mg/ml of NaCl and
 3.0 mg/ml of NagPO, pH7.0.
- INFERGEN® (interferon alfacon-1), made by Amgen Inc., also 25 known as consensus interferon, is marketed for the treatment of adults with chronic Hepatitis C virus (HCV) infections. It is currently the only FDA approved, biooptimized interferon developed through rational drug design and the only interferon with data in the label specifically for non-responding or refractory patients. InterMune's sales force re-launched Infergen® in January 2002 with an active campaign to educate U.S. hepatologists about the safe and appropriate use of Infergen®, which represents new hope for the more than 50 percent of HCV patients who fail other currently available therapies. See

http://www.intermune.com/wt/itmn/infergen, 8/27/2003

Figure 6-B. Circular Dichroism spectrum of Infergen® From Reference [Journal of Interferon and Cytokine Research. 16:489-499(1996)]

Figure 6-C. Circular Dichroism spectrum of rSIFN-co <u>Spectrum range</u>: 320nm-250nm <u>Sensitivity</u>: 2 m*/cm

10 Light path: 2cm

Equipment: Circular Dichroism J-500C

Samples: contains 0.5mg/ml rSIFN-co, 5.9 mg/ml of NaCl and 3.8 mg/ml of Na₂PO₄, pH7.0.

15 Figure 6-D. Circular Dichroism spectrum of rSIFN-co Spectrum range: 250nm - 190nm Sensitivity: 2 m*/cm Light path: 0.20 cm Equipment: Circular Dichroism J-500C

20 Samples: contains 30pg/ml rSIFN-co, 5.9 mg/ml of NaCl and 3.8 mg/ml of Na2PO, pH7.0.

Clearly, as evidenced by the above spectra, the secondary or even tertiary structure of rSIFN-co is different from 25 Inferoen.

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DETAILED DESCRIPTION OF THE INVENTION

This invention provides a recombinant super-compound interferon or an equivalent thereof with changed spatial configuration. This invention reveals that protein with same primary sequence might have different biological activities. As illustrated in the following example, this invention disclosed two proteins with identical smino acid sequence but with different activities. This activity may sometimes become improved efficacy and sometimes, the protein with changed spatial configuration would reveal new function.

An equivalent is a molecule which is similar in function to the compound interferon. An equivalent could be a deletion, substitution, or replacement mutent of the original sequence. Alternatively, it is also the intention of this invention to cover mimics of the recombinant supercompound interferon. Mimics could be a peptide, polyopetide or a small chemical entity.

The interferon described herein includes but is not limited to interferon a, 3, or a. In an embodiment, it is IFN-la, TFN-2b or other mutants.

In an embodiment, the super-compound interferon disclosed has higher efficacy than the interferon described in U.S. Patent Nos. 4,695,623 or 4,897,471. This super-compound interferon is believed to have unique secondary or tertiary structure. (See e.g. Figure 6)

The super-compound interferon described herein has special structure change(s) resulting from the changes of its production process.

The above-described super-compound interferon may be

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produced by a high efficiency expression system which uses a special promoter. In an embodiment, the promoter is Page. As it could be easily appreciated by other ordinary skilled artisan, other inducible promoter, such as heat shock promoter, may be used in this invention.

The super-compound interferon may also be produced with its gene as artificially synthesized cDNA with adjustment of its. sequence from the wild-type according to codon preference of E. Coli. Extensive discussion of said codon usage (preference) may be found in U.S. Patent No. 4,695,623. See e.g. column 6, line 41 - column 7, line 35

The above described super-compound interferon possesses anti-viral or anti-tumor activity and therefore useful in preventing and treating viral diseases, tumors or cancers.

The virus diseases include but are not limited to Hepatitis A, Hepatitis B, Hepatitis C, other types of hepatitis, infections caused by Epstein-Barr virus, Cytomegalovirus, herpes simplex viruses, other herpes viruses, papowaviruses, poxviruses, picornaviruses, adenoviruses, rihnoviruses, human T cell leukaemia viruses I, human T cell leukaemia viruses II, or human T cell leukaemia viruses III.

Therefore, this invention provides a method for inhibitive virus replication or virus infected cells by contacting said virus or infected cells with an effective amount of the super-compound interferon or its equivalent. This super-compound interferon is useful in preventing or treating the following cancers or tumors:

Cancer		Basal Cell Carcinoma
	Skin Cancer	Malignant Melanoma
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	Renal cell carcinoma	
-	Liver Cancer	
H	Thyroid Cancer	
1	Rhinopharyngeal Cancer	
		Prostate Cancer
		Tummy Cancer
	Solid Carcinoma	Esophagus Cancer
		Recta Cancer
	•	Pancreas Cancer
		Mammary Cancer
[Ovarian Cancer & Superficial Bladder	
	Cancer	
	Hemangioma	
		Cervical Cancer
	Epidermoid Carcinoma	Non-small Cell Lung Cance
	* 4	Small Cell Lung Cancer
		Glioma
		Acute Leucocythemia
	Leucocythemia	Chronic Leucocythamia
Malignant	Chronic Myelocytic Leukemia	
Hemal	Hairy Cell Leukemia	
Disease	Lymphadenoma	
	Multiple Myeloma	
	Polycythemia Vera	
Others	Kaposi's Sarcoma	<u> </u>

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Accordingly, this invention provides a method for inhibiting tumor or cannor cell growth by contacting the super-compound interferon or its equivalent vith said tumor or cancer cells. In a further embodiment, the super-compound interferon inhibits the DNA duplication and secretion of RBSAg and HBSAg of Repatits 3 Yurus.

This invention also provides an artificial gene codes for the super-compound interferon or its equivalent. It is within the ordinary skill to design an artificial gene. Hany methods for generating nucleotide sequence and other molecular biology techniques have been described previously. See for example, Joseph Sambrook and David W. Russell, Molecular Cloning: A leboratory Manual, December 2000, published by Cold Spring Harbor Laboratory Press.

This invention provides a vector comprising the gene which codes for the super-compound interferon or its equivalent.

- This invention provides an expression system comprising the vector comprising the gene which codes for the supercompound interferon or its equivalent. The cells include but are not limited to prokaryotic or eukaryotic cells.
- 25 This invention also provides a host cell comprising the vector comprising the gene which codes for the supercompound interferon or its equivalent.
- This invention provides a process for production of recombinant super- compound interferon comprising introducing an artificial gene with selected codon proference into an appropriate host, culturing said introduced host in an appropriate condition for the expression of said compound interferon and harvesting the

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The process may comprise extraction of super-compound interferon from fermentation broth, collection of inclusion body, denaturation and renaturation of the harvested protein.

The process may maintain the high afficacy even when the super-compound interferon is used with an agent and in a particular concentration. The process also comprises separation and purification of the super-compound interferon. The process further comprises lyophilization of the purified super-compound interferon. The process comprises production of liquid injection of super-compound interferon.

This invention also provides the produced super-compound interferon by the above processes.

This invention provides a composition comprising the 20 recombinant super-compound interferon or its equivalent and a suitable carrier.

This invention provides a pharmaceutical composition comprising the recombinant super-compound interferon or its equivalent and a pharmaceutically acceptable carrier.

This invention provides a method for treating viral diseases or tumor in a subject comprising administering to the subject an effective amount of the super-compound interferon or its equivalent.

This invention provides the above-described method wherein the viral diseases is Hepatitis A, Hepatitis B, Hepatitis C, other types of hepatitis, infections of viruses caused by Epstein-Barr virus, Cytomegalovirus; herpes simplex viruses, or other type of herpes viruses, papovaviruses,

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poxviruses, picornaviruses, sdenoviruses, rihnoviruses, human T cell leukaemia viruses I, or human T cell leukaemia viruses II, or human T cell leukamia virus III.

This invention provides the above-described method wherein super-compound interferon was administered via oral, vein injection, muscle injection, peritoneal injection, subcutaneous injection, nasal, mucosal administration, by inhelation via an inspirator.

This invention provides the above-described method wherein super-compound interferon was administered following the protocol of injection 9 µg or 15 µg per day, 3 times a week, totally 24 weeks.

It was surprising to find that rSIPN-co, the spatial configuration of which has been changed, is not only a preparation to inhibit the DNA duplication of Hepatitis B, but to inhibit the eccretion of HBsAg and HBsAg.

One objective of this invention is to offer a preparation of rSIFN-co to directly inhibit the DNA duplication of Hepatitis B viruses and the secretion of Heads and HBsAg of Hepatitis B and decrease them to normal levels.

In one of the results of this invention, rSIFN-co was produced with recombinant techniques. On the condition of fixed amino acid sequence, the IFN DNA was redesigned according to the E. Coli. codon usage and then the rSIFN-co gene was artificially synthesized. rSIFN-co CDNA was cloned into the high-expression vector of E. Coli. by DNA recombinant techniques, and a high expression of rSIFN-co was gained by using of induce/activate-mechanism of Larabinose to activate the transcription of Pan promoter.

Compared with usual thermo-induction, pH induction and IFTG

induction systems of genetic engineering, arabinose induction/activation system has some advantages: (1) Common systems relieve promoter function by creating a "derepression" pattern. Promoters then induce downstream dene expression. So temperature and pH change and the addition of IPTG cannot activate promoters directly. In the system disclosed herein, L-arabinose not only deactivates and represses but also activates the transcription of Pano promoter which induces a high expression of rSIFN-co. Therefore, the arabinose induction/activation system is a more effective expression system. (2) The relation between Exogenous and L-arabinose dosage is linearity. This means the concentration of arabinose can be changed to adjust the expression level of the exogenous gene. Therefore, it is easier to control the exogenous gene expression level in £. Coli. by arabinose than by changing temperature and pH value. This characteristic is significant for the formation of inclusion bodies. (3) L- arabinose is resourceful cheap and safe, which, on the contrary, are the disadvantages of other inducers such as IPTG.

This embodiment creates an effective and resistant rSIFNco-expressing E. Coli. engineering strain with an Larabinose induction/activation system. The strain is cultivated and fermented under suitable conditions to harvest the bacterial bodies. Inclusion bodies are then purified after destroying bacteris and washing repeatedly. The end result, mass of high-purity, spaticial-structurechanged rSIFN-co protein for this invention and for clinical treatment, was gained from denaturation and renaturation of inclusion bodies and a series of purification steps.

The following are some rSIFN-co preparations: tablets, 35 capsules, oral liquids, pastes, injections, sprays, suppositories, and solutions. Injections are recommended. It is common to subcutaneously inject or vein-inject the medicine. The medicine carrier could be any acceptance medicine carrier, including carbohydrate, cellulosum, adhesives, disintegration agents, emcilients, filling, addissolve agent, amortization, preservative, add-thick agent, matching, etc.

This invention also provides a pharmaceutical composition comprising the above composition and a pharmaceutically 10 acceptable carrier.

For the purposes of this invention, "pharmaceutically acceptable carriers" means any of the standard pharmacoutical carriers. Examples of suitable carriers are well known in the art and may include, but are not limited to, any of the standard pharmaceutical carriers such as a phosphate buffered saline solution and various vetting agents. Other cerriers may include additives used in tablets, granules and capsules, etc. Typically such carriers contain excipients such as starch, mik, sugar, certain types of clay, gelatin, stearic acid or salts thereof, magnesium or calcium stearate, talc, vegetable fate or oils, gum, glycols or other known excipients. Such carriers may also include flavor and color additives or other ingredients. Compositions comprising such cerriers are formulated by vell-known conventional methods.

This invention will be better understood from the examples which follow. However, one skilled in the art will readily appraciate that the specific methods and results discussed are merely illustrative of the invention as described more fully in the claims which follow thereafter.

EXPERIMENTAL DETAILS

EXAMPLE 1

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rSIFN-co is a new interferon molecule constructed according to conservative amino acid in human IFN-c subtype with questic engineering method. It has been proven that rSIFNco has broad-spectrum IFN activity, such as high antivirus and tumor inhibition activity, especially for effectively treating legativis C.

E. Coli. codon was used to redesign rSIFN-co cDNA and then artificially synthesize cDNA of rSIFN-co from published rSIFN-co DNA sequences and deduced amino acid sequences (Figure 1).

In order to get pure rSIFN-co protein, rSIFN-co CDNA was cloned into E. Coli. high-expression vector, and L-arabinose, which can activate strong PBAD promoter in vectors, was used to induce high expression of rSIFN-co cere.

- 1. Synthesis of E. Coli. cDNA Sequence
- 1.1 Redesign of rSIFN-co cDNA sequence
- or FSIEN-co oDNA was redesigned according to the codon usage of E. Coli. to achieve high expression in E. Coll. Deduced amino soid sequence from the redesigned cONA sequence of rSIEN-co is completely coincidental with primitive amino acid sequence of published rSIEN-co (Figure 1).
- 25 1.2 rSIFN-co cDNA sequence synthesis
 - 1.2.1r9IFN-co cDNA 5'-terminus and 3'- terminus semimolecular synthesis

Two semi-moleculars can be directly synthesized: rSIFN-co CDNA 5'- terminus 280bp (fragment I) and 3'- terminus 268bp(fragment II) by PCR. There are 41bp overlapping among fragment II and fragment I.

- (1) Chemical synthesis oligodeoxynucleotide fragment:
- Oligomer A:
- 35 5'ATCTGCGACCTGCCGAGACCCACTCCCTGGGTAACCGTCGTGCTCTGTTCTGCTCG GATGCGTCGTATCTCCCGTTCTCCTGCCTGAAAAACCCGTCACGAC3'

Oligomer B:

 ${\tt 5'CTGAAAGACCGTCACGACTTCGGTTTCCCGCAGGAGAGGTTCGACGGTAACCAGTTCCAGA}$ AGCTCAGGCTATCTCCGTTCTGCACGAAATGATCCAGCAGACCTTC3'

Oligomer C:

5'GCTGCTGGTACAGTTCGGTGTAGAATTTTTCCAGCAGGGATTCGTCCCAAGCAGCGGAGGAG TCTTTGGTGGAGAACAGGTTGAAGGTCTGCTGGATCATTTC3'

Oligomer D:

5'ATCCCTGCTGGAAAATTCTACACCGAACTGTACCAGCAGCTGAACGACCTGGAAGCTTGCG TTATCCAGGAAGTTGGTGTTGAAGAACCCCGCTGATGAAC3'

Oligomer E:

5 GAAGAAACCCCGCTGATGAACGTTGACTCCATCCTGGCTGTTAAAAATACTTCCAGCGTAT CACCCTGTACCTGACCGAAAAAAAATACTCCCCGTGCGCTTGGG3'

Oligomer F:

PCR I for Fragment I: oligodecxynucleotide B as template, oligodecxynucleotide A and C as primers, synthesized 280 bp Fragment I.

20 PCR I mixture (units: ul)

sterilized distilled water	39
10×Pfu buffer (Stratagen American Ltd.	5
dNTP mixture (dNTP concentration 2.5 mmol/L)	2
Oligomer A primer (25 µmol/L)	1
Oligomer C primar (25 µmol/L)	1
Oligomer B template (1 µmol/L)	1
Pfu DNA polymerase (Stratagen American Ltd.) (25 U/µ1)	1
Total volume	50ul

PCR cycle: 95 I

2m-(95°C45s-65°C1m-72°C1m) ×25 cycle-72°C10m-4°C

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PCR II for Fragment II: oligodeoxynucleotide E as template, oligodeoxynuclectide D and F as primers, synthesized 268bp Fragment II. PCR II mixture (units: pl)

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sterilized distilled water	39
10*Pfu buffer (Stratagen American Ltd.)	5
dNTP mixture (dNTP concentration 2.5mmol/L)	2
Oligomer D primer (25 µmol/L)	1
Oligomer F primer (25 µmol/L)	1
Oligomer E template (1 µmol/L)	1
Pfu DNA polymerase (Stratagen American Ltd.)	(25U/µl) 1
Total volume	50µ1
PCR cycle: the same as PCR I	

1.2.2Assembling of rSIFN-co cDNA

Fragment I and II were assembled together to get the complete cDNA molecular sequence of rSIFN-co using the overlapping and extending PCR method. Restriction snzyme Nde I and Pst I were introduced to clone rSIFN-co cDNA sequence into plasmid.

(1) Chemical synthesis primers
Oligomer G: 5'ATCGGCCATATGTGCGACCTGCCGCAGACCC3'

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Oligomer H: 5'ACTGCCAGGCTGCAGTTATTCTTTACGACGCAGACGTTCC3'
(2) Overlapping and extending PCR

PCR mixture (units:	μl)
sterilized distilled water	38
10×Pfu buffer (Stratagen American Ltd.)	5
dNTP mixture (dNTP concentration 2.5mmol/L)	2
primer G (25 µmol/L)	1
primer R (25 µmol/L)	1
*fragment I preduction (1 µmol/L)	1
*fragment II preduction (1 umol/L)	1
Pfu DNA polymerase (Stratagen American Ltd.) (2.50/ μ l)	1
Total volume	50u

*Separate and purify PCR production with StrataPrep PCR purification kit produced by Stratagen American Ltd. And dissolve into sterilized distilled water.

Leu-

PCR cycle: the same as PCR 1

2. rSIFN-co gene clone and sequence analysis
plac T7 plasmid as cloning vector. plac T7 plasmid is

reconstructed with pBluescript II KS(+) plasmid produced by Stretagen (Figure 3).

Purified PCR production of rSIFN-co CDNA with StrateProp PCR purification kit. Digest CDNA and plac T7 plasmid with Medr and PstI. Run 18 agarose gel electrophoresis and separate these double-digested DNA fragments. Recover 507bp long rSIFN-co DNA fragment and 2.9kb plasmid DNA fragment. Ligate these fragments by T4 DNA ligase to form a recombinant plasmid. Transform DH_{Sc}Competent cells (Gibco) with the recombinant plasmid. culture at 37°C overnight.

Identify the positive recombinant colony, named pHY-1.

Run DNA sequencing with SequithhermIM Cycle Sequencing Kit produced by American Epicentre Technologies Ltd using Li-COR Model 400UL. Primers are T7 and T3 common sequence primer, the DNA sequencing result matches theoretic design.Purify the rSIFM-co, sequence the N-terminus amino acids. The N-terminus amino acids sequence matches

experimental design which is as follows:

N- Cys-Asp-Leu-Pro-Gln-Thr-His-Ser-Leu-Gly-Asn-Arg-Arg-Ala-

- 25 3. Construction, transformation, identification, and hereditary stability of expression vector
 - 3.1 Construction and transformation of expression vector

Digested E. Coli. expression vector pHY-4(see Figure 3), with Nde I to linearize and subsequently digest with Xba I.

30 Run 1% agarose gel electrophoresis, and purify the 4.8kb pHY-4 Nde I -Xba I digest fragment with QIAEX II kit produced

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by QIAGEN Germany Ltd.

At the same time, the pHY-4 plasmid is double digested with Nde I-Xba I. Run lt agarose gel electrophoresis and purify the 715bp fragment. Ligate the rSiFN-co and pHY-4 fragments with T4 DNA ligase to construct the recombinant plasmid (See Figure 4). Transform OH₃₀competent cells with the recombinant plasmid. Spread the transformed cells on LB plate with Amp. 37°C culture overnight.

3.2 Positive closing strain screening

- Randomly choose E. Coli. colonies from above LB-plate, screening the positive strains containing recombinant vector by endonuclease digesting and PCS analysis. Name one of the positive recombinant plasmid pHY-5, and name the strain containing pHY-5 plasmid PVIII. Amplify and store the positive strain with qluverol in -80°C.
- 4. High expression of rSIFN-co gene in E. Coli.

 In pRY-5 plasmid, rSIFN-co gene is under control of strong promoter Page. This promoter is positively and negatively regulated by the product of the gene arc. AraC is a transcriptional regulator that forms a complex with arzbinose. In the absence of argbinose, the AraC dimer blinds O2 and I. forming a 210bp loop. This conformation leads to a complete inhibition of transcription. In the presence of arabinose, the dimer is released from O2 and binds I1 and I2 leading to transcription. Arabinose binding desctivates, represses and even activates the transcription of Page promoter, which stimulates Page inducing high expression of rSIFN-co. rSIFN-co expression level in PVIII is more than 50% of the total E. Coli. proteins.
 - 5. Summary rSIFN-CO is a new interferon molecule artificially built

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according to the conservative smino acid of human α interferons. It has been proven as a effective antihepatitis drug. In order to get enough pure rSIFN-co protein, a stable recombinant E. Coli. strain which high expresses rSIFN-co protein was constructed.

First, according to published rSIFN-co amino acid sequence, E. Coli. codon was used to synthesize whole CDNA of RSIFN-co. This DNA fragment was sequenced and proved that the SOlbp codon sequence and TAA termination codon sequence are valid and identical with theocratic design. Subsequent analysis revealed that the N-terminus amino acid sequence and amino acid composed of rSIFN-co produced by the recombinant strain were both identical to the prediction.

The rSIFN-co cDNA was cloned into E. Coli. high-expression vector pHY-4 plasmid to construct the recombinant plasmid pHY-5. E. Coli. LMG194 strain was further transformed with pHY-4 plasmid to get stable rSIFN-co high-expression transformant. This transformant was cultured for 30 generations. The heredity of pHY-5 recombinant plasmid in E. Coli. LMG194 was normal and stable, and the expression of rSIFN-co was high and steady.

25 E. Coli. LMG194, which contains recombinant pHY-5 plasmid, is actually an ideal high-expression engineering strain.

6. References

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30 EXAMPLE 2

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Separation and purification of rSIFN-co

1. Fermentation

Inoculate the recombinant strain in LB media, shaking (200 rpm) under 37°C overnight (approximate 18 h), then add 30%

glycerol to the fermentation broth to get final concentration of 15%, allotted to 1 ml tube and kept in - $20\,^{\circ}\text{C}$ as seed for production.

5 Add 1% of the seed to LB media, shaking (200 rpm) under 37°C overnight to enlarge the scale of the seed, then add to RM media with a ratio of 105, culturing under 37°C. Add arabinose (20% solution) to 0.02% as an inductor when the OD600 reaches about 2.0. 4 hours after that, stop the culture process, collect the bacteria by centrifuge, resuggend the pellet with buffer A, and keep in -20°C overnight. Thaw and break the bacteria by homogenizer, then centrifuge. Wealt the pellet with buffer B, buffer C, and distilled water to get a relatively pure inclusion body.

2. Denaturation and renaturation

Dissolve the inclusion body in Guanidine-RC1 (or ures) of 6 mol/L. The solution will be a little cloudy. Centrifuge it at a speed of 10000 rpm. Determine the protein concentration of the supernatant. This supernatant is called "denaturation solution." Add the denaturation solution to renaturation buffer, and keep the final protein concentration under 0.3 mg/ml. It is better to add the totally denaturation solution in three steps instead of one step. Keep the solution overnight under 4°T. Afterwards, dialyze against 10 mol/L and 5 mol/L PB buffer and distilled water, then adjusting its pib by 2 mol/L RAc-NaAc. Let it standstill for a while, then filtrate.

30 3. Purification

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POROS HS/M anion exchange chromatography: Equivalent column with 20 mmol/L HAc-NaAc(pH 5.0)

Load samples at a speed of 30 ml/min

22

30

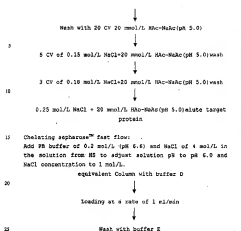
Wash with 20 CV 20 mmol/L HAC-NaAc(pH 5.0) 5 CV of 0.15 mol/L NaCl+20 mmol/L HAc-NaAc(pH 5.0) wash 3 CV of 0.18 mol/L NaCl+20 mmol/L HAc-NaAc(pH 5.0) wash 0.25 mol/L NaC1 + 20 mmol/L HAc-NaAc(pH 5.0) elute target protein Chelating sepharose™ fast flow: Add PB buffer of 0.2 mol/L (pH 6.6) and NaCl of 4 mol/L in the solution from HS to adjust solution pH to pH 6.0 and NaCl concentration to 1 mol/L. equivalent Column with buffer D Loading at a rate of 1 ml/min Wash with buffer E Wash with buffer F

Condense the eluted solution by POROS H5/M. Sometimes a step of purification by sephacryl S-100 can be added to meet with stricter purity requirements.

2

Elute with buffer G

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Condense the eluted solution by POROS H5/M. Sometimes a step of purification by sephacryl S-100 can be added to meet with stricter purity requirements.

23

Wash with buffer F

Elute with buffer G

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Note:

Buffer A: 100 mmol/L Tris-HCl,pH 7.5-10 mmol/L EDTA-100 mmol/L NaCl

- Buffer B: 50 mmol/L Tris-HCl,pH 7.5-1 mol/L Grea-10 mmol/L EDTA-0.5% Triton X-100 Buffer C: 50 mmol/L Tris-HCl,pH 7.5-2 mol/L Urea-10 mmol/L
 - EDTA-0.5% Triton X-100 Buffer D: 1 mol/L NaCl ---50 mmol/L Na2HPO4 (pH 5.5)
- Buffer E: 1 mol/L NaCl ---50 mmol/L Na2HPO4 (pH 5.0) 10
 - Buffer F: 1 mol/L NaCl --- 50 mmol/L NasHPO4 (pH 4.0)

Buffer G: 1 mol/L NaCl ---50 mmol/L Na-HPO, (pH 3.6) Renaturation buffer: 0.5 mol/L Arginine-150 mmol/L Tris-HCl. pH 7.5-0.2 mmol/L EDTA

- LB Media: 1 L 15
 - Tryptone 10 a
 - Yeast extracts 5 q N=C1 10 a
 - RM Media: 1 L
- Casein 20 a
 - 1 mmo1/L (0.203 g) MgC1
 - Na₂HPO₄ 4 g;
 - KH2PO4 3 0,
 - NaCl 0.5 a
- 25 NH₄C1 1 g

After purification, the buffer was changed to PBS (pH 7.0) along with the step of condensing by POROS HS/M. This is called the "Protein Stock Solution." It can be directly used in the preparation of injections or sprays, or stored at 2-8 degree centigrade.

Formula for injection:

Solution of rSIFN-	Solution 34.5 µg/ml	Lyophilized powder 34.5 µg/ml
co FB (pH7.0) Glycine NaCl	25mmo1/L 	10mmol/L 0.4mol/L
For spray: EDTA Tween 83 Trisodium citrate Glycerol Sodium Chloride Phenylmethanol HSA rSIFN-co	0.01% 0.05% 10mmol/L 1.26% 0.03% 0.5% 0.1% 10 µg/ml	*

5 QUALITY CONTROL PROCESS

During purification tests for protein content, protein purity, specific activity and pyrogen are conducted after each step. When the stock solution is obtained, all the tests listed in the table are done one after the other.

The quality of the product is controlled according to "Chinese Requirements for Biologics"

1. Original protein solution

Lowry

10

Item of Test	Method
Protein Stock Solution:	
lest for Protein Content	Lowry
Test for Protein Purity	Non-reductive SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) MPLC Analysis
lest for Molecular Weights	Reductive SDS-FAGE

2:

777 - 846 - 177	According to Method in
Test for Specific Activity	"Specific Activity Test of
	Interferon
	Using DNA Labeling and
Test for Leftover Exogenetic	Detection Kit
DNA	According to Method in
Test for Activity of	"Chemical and Other Test
Leftover Antibiotics	Methods for Biologics"
Test for Bacterial Endotoxin	"Requirements for Bacterial
	Endotoxin Test of Biologics"
	Isoelectric Focusing
Test for Isoelectronic Point	Isoelectric rocusing
	Electrophoresis
Test for Identify	UV spectrum (range of
Characteristics of the	wavelength: 190-380nm)
Protein	
	Peptide Mapping (hydrolyzed
	by pancreatic enzyme,
	analyzed by C-16 column)
	N-terminal Sequence Test
	C-terminal Sequence Test
	Circular Dichroism
	Amino Acid Analysis
Semi-finished Product	
Test for Bacterial Endotoxin	According to Method in
1050 101 501101	"Requirements for Bacterial
	Endotoxin Test of Biologics"
Product	
Appearance Check	7
Chemical	According to Method in
Chemicai	"Chemical and Other Test
	Methods for Biologics"
Test for Specific Activity	According to Method in
Jeac for observer wertarn	"Specific Activity Test of
	Interferon
Sterility Test	According to Method in "c"
Abnormal Toxicity Test	Test on Mouse
Abnormal Toxicity Test	According to Method in
Pyrogen Test	"Requirements for Pyrogen
	Test of Biologics"
	Test of providing
Test for Stability of	1
Product	

Note: "Chemical and Other Test Methods for Biologics",

"Requirements for Pyrogen Test of Biologics" and

"Requirements for Bacterial Endotoxin Test of Biologics",

all can be found in the "Chinese Requirements for

Biologics." "Chinese Requirements for Biologics," FAN

Zhengan, ZHANG Xinhui, DUAN Zhibing, et al. Chinese Biologics Standardization committee. Published by Chemical Industry Publishing Company, 2000.

5 EXAMPLE 3

Stability of lyophilized Fowder of Recombinant Super-Compound Interferon Injection

The stability experiments were carried out with samples of 10 lyophilited powder of recombinant super-compound interferon (rSIFN-co) injection in two specifications and three batches. The experiments started on April, 2000.

1. Sample Source

15 Samples were supplied by Sichuan Huiyang Life-engineering Ltd., Sichuan Province. Lot: 990101-03, 990101-05, 990102-03, 990102-05, 990103-03, 990103-05

2. Sample Specifications

20 Every sample in this experiment should conform with the requirements in the table below.

Table 1 Standards of Samples in Experiment

Items	Standards
1. Appearance	white loose powder
2. Dissolving	dissolve rapidly in injection water(within
time	2 min) at room temperature
3. Clarity	colourless liquid or with little milk-like glisten: should not be cloudy, impurity or with indiscerptible deposit
4. pH value	6.5~7.5
5. Potency (IU/dose)	809-150% of indicated quantity (9µg:4.5 × 10°IU, 15µg: 7.5 × 10°IU)
6. Moisture	no more than 3.0% (W/W)

3. Experiment Content

2

15.3.1 Test samples at 2-8°C: The test samples were put into a 2-8°C refrigerator, then the above items of 6°f sea samples were respectively tested in the $1^{\rm Sc}$, $3^{\rm Sd}$, $6^{\rm Sh}$, $3^{\rm Sh}$, $26^{\rm Sh}$, $36^{\rm Sh}$ month. The results were recorded.

15.3.2 Test samples at 25°C: The test samples were put into a thermostat at 25°C, then the above items of these samples were respectively tested in the 1st., 3rd., 6ts., 9tm., 12°th, 19th, 20°M month. The results were recorded.

15.3.3 Test samples at 37°C: The test samples were put into a thermostat at 37°C, then the above items of these samples were respectively tested in the 1°°. 3°d. 6°n. 9°h. 12°n. 13°b. 24°n month. The results were recorded.

4. Results and Conclusion

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1) At 37°C, according to data collected at designated points during testing and compared with data before testing, the potency began descending from the 6th month and the changes in the three batches were similar. The supearance of other items had no changes.

2) At 25°C, according to data collected at designated points during testing and compared with data before the testing, the potency only had a little change, and the changes in the three batches were similar. The appearance of other trems had no changes.

 At 2-8°C, according to data collected at designated points during testing and compared with data before testing, the potency of the three batches all were stable. The appearance of other items also had no changes.

In conclusion, it is suggested that the lyophilized powder of recombinant super-compound interferon for injection should be better stored and transported at low temperatures. Without such conditions, the product can also be stored for short periods (i.e. 3 months) at room

temperature.

EXAMPLE 4

rSIFM-co inhibits HBV-DNA duplication and secretion of HBsAg and HBsAg.

Materials

Solvent and Dispensing Method: Add lml saline into each vial, dissolve, and mix with MEM culture medium at different concentrations. Mix on the spot.

Control drugs: IFN-a2b (Intron A) as lyophilized powder, purchased from Schering Flough. 3×10⁴U each, mix to 3×10⁴U/ml with culture medium; INFERGEM⁹ (liquid colution), purchased from Amgen, 9µg. 0.3ml each, equal to 9X10⁴U, and mix with 9X10⁴U/ml culture medium preserve at 4°C; 2.2.15 cell: 2.2.15 cell line of hepatoms (Hep G2) closed and transfected by HBV ONA, constructed by Mount Sinai Medical Center.

Reagent: MEM powder, Gibco American Ltd. cattle fetal blood gerum, HycloneLab American Ltd. G-418(Geneticin); MEM dispensing, Gibco American Ltd.; L-Glutemyl, imported and psokaged by JING KE Chemical Ltd.; HBsAg and HBsAg solidphase radioimmunossay box, Northward Reagent Institute of Chinese Isotope Ltd.; Biograncetina, Northern China Medicine, And Libofectin, Gibco American Ltd.

Experimental goods and equipment: culture bottle, Denmark
O Tunclon^N, 24-well and 95-well culture board, Corning
American Ltd.; Carbon Dioxide hatching box, Shal-Lab
American Ltd.; MEM culture medium 100ml: 10% cattle fetal
blood serum, 3% Glutamyll%, G418 380ug/ml,
bloodranectima500/ml.

35 Method:

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2.2.15 cell culture: Added 0.25t pancreatic enzyme into culture box with full of 2.2.15 cell, digest at 37°C for 3 minutes, and add culture medium to stop digest and disturb it to disperse the cells, reproduce with ratio of 1:3. They will reach full growth in 10 days.

Toxicity test: Set groups of different concentrations and a control group in which cell is not acted on with medicine. Digast cell, and dispense to a 100,000 cell/ml solution. Inoculate to 96-well culture board, 200ml each well, culture at 37% for 24h with 5% CO2. Test when simple cell layer grows.

Dispense rSIFN-co to 1.8×10'IU/ml solution than prepare a series of solutions diluted at two-fold gradients. Add into 96-well culture board, 3 wells per concentration. Change the solution every 4 days. Test cytopethic effect by microscope after 8 days. Fully destroy as 4, 75% as 3, 50% as 2, 25% as 1, zero as 0. Calculate average cell lesion and inhibition rate of different concentrations. Calculate 7550 and TCO according to the Reed Muench method.

TC50 = Antilog (B +
$$\frac{50-B}{A-B}$$
 × C)

A=log >50% medicine concentration, B=log<50% medicine concentration, C=log dilution power

Inhibition test for HBeAg and HBsAg: Separate into positive and negative HBeAg and HBsAg contrast groups, cell contrast groups. The deficine concentration groups. Inoculate 700,000 cells/ml of 2.2.15 cell into 6-well culture board, 3 ml each well, culture at 37°C for 24h with 55 CO₂, then prepare 5 gradiently diluted solutions with 3-fold as the grade (Prepare 5 solutions, each with a different protein concentration. The concentration of Solution 2 is 3 times lower than that of Solution 1, the concentration of solution 3 is 3 times lower than that of Solution 2, etc..)

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4.5*10*TO/ml, 1.5*10*TO/ml, 0.5*10*TU/ml, 0.17*10*1U/ml, and 0.056*10*TU/ml, 1 well per concentration, culture at 37°C for 24h with 58 CQ. Change solutions every 4 days using the same solution. Collect all culture medium on the 8° day. Preserve at -20°C Repeat test 3 times to estimate HBsAg and HBsAg with solid-phase radicimaunossasy box (Northward Reagent Institute of Chinese Isotope Ltd.). Estimate comvalue of each well with a y- accounting machine.

10 Effects calculation: Calculate cpm mean value of contrast groups and different-concentration groups and their standard deviation, P/N value such as inhibition rate, IC50 and SI.

1) Antigen inhibition rate (%) = $\frac{A-B}{A} \times 100$

A - com of control group; B - com of test group;

2) Counting the half-efficiency concentration of the medicine

Antigen inhibition IC50 - Antilog (B + $\frac{50-B}{A-B}$ x C)

A-log-504 medicine concentration, B-log<504medicine
concentration, C-log dilution power

 SI of interspace-conformation changed rSIFN-co effect on HBeAg and HBeAg in 2.2.15 cell culture:

SI - TC50

4) Estimate the differences in cpm of each dilution degree from the control group using student t test

Southern blot: (1) HBV-DNA extract in 2.2.15 cell: Culture cell 8 days. Exauction culture medium (Separate cells from culture medium by means of draining the culture medium.). Add lysis buffer to break cells, then extract 2 times with a mixture of phanol, chloroform and isoamyl aloohol

31

(1:1:1), 10,000g centrifuge. Collect the supernatant adding anhydrous alcohol to deposit nucleic acid. Vacuum draw, redissolve into 20piTE buffer. (2) Electrophoresis: Add 6XDNA loading buffer, electrophoresis on 1.5% agarose gel, IV/cm, at fixed pressure for 14-18h. (3) Denaturation and hybridization: respectively dip gel into HCI, denaturation buffer and neutralization buffer. (4) Transambrane: Make an orderly transfer of DNA to Hybond-N membrane. Bake, hybridize and expose with dot blot hybridization. Scan and analyze relative denaity with gel-pro software. Calculate inhibition rate and ICSO.

Results

Table 1: Results of inhibition rate of rSIPN-co to HBsAg and HBeAg Wiset hanch. FREEN-col

				-	Inhibition effect to HBeAg	effect to	HBeAg			
	Pire	Secon	Thir	Linki	Inhibition rate		Avetage	Accusulation	1	Accumulated
n(x10'IU/ml)	t well	Į,	d 1194	First	Second	Third well	inhibitio n rate	a	Accumulation	rate
906	9026	8976	1043	0.436227	0,43935	9.34565	0.407079	0.945909	0.592921	0.614693546
300	9616	12082	1008	0.399375	9.24534	9.36926	0.337997	0.5388299	1.254924	0.300392321
. 001	9822	16002	1280	0.366508	0.0005	0.2005	0.195836	0.200833	2.059088	0.00867188
33.33333	1577	19306	1682	0.014991	0	0	166900.0	0.0049969	3.054091	0.601633453
111111	1917	22270	1893	0	0	0	0	0	4.054091	0
Control	100	16010		Blank	۰		Dilution	3	1050	6
			Inh	Inhibition effect to aBakg	act to HBs	SV1				
O Jaco descend	Firs	Secon	Thir	Any	Inhibition rate	te	Average	Accumulatio	1-	Accountated
(*10,10,m)	¥ 1	4 MB21	d #eII	First	Second	Third	n rate	a	n a	rate
200	7706	7240	13.4	0.342155	0.38193 6	0.39269	D.372261	0.922258	0.627739	0.595006426
300	9886	8777	9476	0.243981	0.33600	0.19105 3	0.257014	0.5499972	1.370724	0.286349225
001	1081	10720	1033	0.07649	0,08485	9.11614	0.093165	0.292983	2.27756	0.113977019
33.3333	če.	ALLI	1057	0.082807	0.05122		0.07723	0.1998179	3.20033	0.058767408
11.1111	1067	9352	1081	0.088953	9,20163	3.07717	0.122588	0.122588	4.077742	0.02918543
Control	5	11714		Blank	0		Dilution	т	1050	641.7736749

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Second batch: [rSIFN-co)

					Tuhib	Ltion affe	Inhibition affact to HBeAq			
	12	Secon	Thir	451	Inhibition rate		Average		-1	Accountate
n(*10'fU/ml)	1 H	rel]	real!	Flrst well	Second	Third well	inhibitio p rate	T.	Accumulation n	inhibition
906	1818	8516	9350	0.554378	0.51459	0.46785	0.512008	1,371181	0,467992	6.73752197
300	103	10628	9160	9,410396	0.39420	6.47788	0.427497	0.8591731	1.060496	0.44756324
100	1229	14228	1326	0.299134	0.18901	0.24407	0.244072	0.4316522	1.816423	0.19201839
33.33333	1536	17414	1618 B	0.124259		0.77291	0.069653	0.1876045	2,74677	0.06393338
11.11131	1738	13632	1540	9.00500.0	0.22298	0.12186 5	0.117951	0.117951	3.628819	0.03148073
Control	Ge LI	1,6962		Blank			Dilution	3	0521	365, 935784 6
			Inh	Inhibition off	offect to HBsAq	Ş.				i
	Fire	Secon	This	Light	Inhibition rate	to.	Average	-,,	-1	Accommiste
n(*10'TU/ml)	rell vell	d well	Į,	First	Second	Third well	inhibitio n <i>r</i> ate	n n	Accumulatio n	inhibition
006	5784	6198	5732	0.498265	0.46235	0.49757	0.486063	0.893477	0.513937	0.63483584
300	7250	8534	8318	177978.0	0.25971 5	6.27845	0.30596	0.4074138	1.207957	0.25221064
100	9830	11212	1021	G.147294	0.02741	0.11433	0.096345	0.101434	2.111612	0.04583464
33.33333	1394	12368	1347	o	0	0		0.0050891	3.111612	0.00163283 5
11.11131	1241 8	11634	1135	0	0	0.01526	69050010	0.005089	4.106523	0.00123772 8
Control	TT Se TT			Blank	۰		Dilution		1050	611.091956 8

¥

Third batch: (rSlFN-co)

					Inhibit	tion effec	Inhibition effect to HBeAg			
Concentration(x	Phrat	Second	This red	HEI.	Inhibition rate	te.	Average	Appendict of	<u>.</u>	Accumulated
10*IU/ml)	T T	1104	We21	First	Second	Third	inhibitio n rate	n accumulation	Accumulat 100	inhibition
900	9702	\$195	8110	0.428016	0.433204	0.52167	0.461031	1,316983	0.538969	0.709599543
300	8314	10032	9870	0.4744723	0.40856	0.47706 6	0.453366	0.8559525	1.085603	0.440859127
100	16312	12688 -	13934	0.038321	0.251975	0.17851	0.156271	0.402586	1.929332	0.172641621
33, 33333	15080	12874	13288	0.110954	0.246547	0.21660	0.190701	0.2463153	2,738631	0.082519158
11.11111	21926	15366	15728	0	0.094093	0.07275	0.0055615	0.055615	3.603017	0.014875633
Control	Ce11	17544		Blank			Dilution	3	ICSD	382.0496935
			Inhibi	Inhibition effect to MasAg	to XBsAg					
Concentration(*	First	Second	This rd	Ioh	Inhibition rate	5	Average	amount to the	À	Accumulated
10'TU/e1)	Hell	¥e11	1	First	Second	Third	inhibitio n rate	n n	Accomulation	inhibition
006	9199	6228	5346	0.496864	0.442035	0.52105	0.486651	0.763125	0.513349	0.597838293
300	8542	8590	3607	0.234725	0.230425	0.35427	0.276474	0.2764738	1.236875	0.182690031
007	11420	11360	11394	0		0	0	0	2.236875	0
33.33333	12656	115#2	13110	0			0	0		
11.1111	13142	12336	13342	0	۰	•	0	0	4.236875	
Control	Ce11	11528		Blank			Dilution	3	ICSO	694, 7027149
Whole Average 1050.		TEVE UST	·uo	139 315476						

Weeleg: Average ICSO: 450.2434 SD: 132.315479 Mosley: Average ICSO: 649.1694 SD: 42.29580 ×

Table 2: Results of inhibition rate of introm A(IEN-a2b) to Healy and HBeAq

						Inhibitio	Inhibition effect to HBeAg	Beng		
Concentration	11.54	Second	1	Thu	Inhibition rate	ate	Average			Accumulated
(*10'TU/ml)	¥ J	Fee13	well	Sirst Well	Second well	Third well	inhibition rate	Accumulation	Accumulation	inhibition rate
300	14918	11724	9950	0	0.029711	0.176529	0.068747	0.068747	0.931253	0.068746724
901	14868	16890	15182	۰	0	0	0	. 0	1.931253	0
33.3333	16760	21716	16400		٥	٥	0	P	2.931253	0
11.1111	75802	15042	16168	0	٥	0	0	0	3.931253	
3.703704	12083	12083	12083	·	0	0	0	0	4.931253	
Control	Ce) I	17544		Blank	۰		Dilution	3	1050	FALSE
			Empi	Inhibition offect to HBsAg	fect to MB	sAg				
Concentration	First	Second	Phi nd	Inh	Inhibition rate	ate	Average			Accumulated
(*10/11/01)		well		First	Second	Third	inhibition rate	inhibition Accumulation rate	Accumulation	inhibition rate
300	9226	9618	9658	0.152489			0.1708	0.189295	0.8292	0.185857736
100	10946	10340	10828	0	0.050156		0.364272 0.018495	0.0184947	1.810705	0.010110817
33,33333	12250	12980	13934	۰	0	۰	0	0	2.810705	0
נזננו.נו	12634	12342	12000	0	0	۰		0	3.816705	0
3.703704	10886	16886	98801		0	0	0	0	4.810705	0
Control .	Ce11	10386		Blank	0		Dilution	3	rc50	PALSE

Table 3: Results of inhibition rate of Infergen[®] to KB4Ag and HBeAg First batch: (Infergen[®])

						-				
						Inhibition	Inhibition effect to Abeng	Deag		
	Firs	Secon	mir	Inh	Inhibition rate	te	Avetage	Beenmy at to	1	Accumulate
Concentration n(*10*10/ml)	1199	d well		First Well	Second	Third well		n	Accumulatio	inhibition
906	2 2 2	12156	_	0.091655	9.22086		0.104175	0.306157	0.895825	9
300	1339	12288	1625	0.141776	9.21240	·	0.118062	0.2019827	1.777764	9
100	1436	18834	1419	0.079349	9	0.09024 5	0.056531	0.083921	2.721232	0.02991667
33.3333	1572	16034	1634	۰	0	0	0	0.0273897	3,721232	0.00730659
1111111	1750	17652	1432	0	٥	0.08216 9	0.02739	0.02739	4.693843	9.00580137
Control	Sell	15602		Blank			Dilution	3	1050	FALSE
			di Hari	Inhibition effect to Ready	ect to HB:	PAG.				
	FARS	Secon	Thir	Tah.	Inhibition rate	te	Average	Accused at io	7	Accomulate
n(*10'IU/nl)	t well	d well	Fe II	First	Second	Third well	ionibitio n rate	a	Accumulation	inhibition rate
006	8021	11692	1223		0.01235	,0	0.00425	0.025163	0.99575	0.02464711
300	1284	11484	1235	۰	0.03031	5	0.010104	0.0209125	1.985646	3 01042207
100	1289	14696	1508	٠		o	o	0.010808	2.985646	0.00360695
33.3333	1503	12928	1302		۰	٥	o	1808010.0	3.985646	6,00270441
11.1111	1179	11984	1150	0.004137	0	0.02828	808010.0	D.010808	4.974837	0.00216783
Control	118	11843		Blank			Dilution	3	1050	FALSE

Table 3: Results of inhibition rate of Infergen* to HEshg and SEchg Error batch: (Infergen*)

riest parcu: (interdeu)	Brank								-	
						Inhibition	Inhibition affect to AbeAg	BeAg		
	Fire	Secon	Thir	Inh	Inhibition rate	9	Average	Possessi at to	4	Accumulate
n(*10*10/ml)	. well	well well	Well d	First	Second	Third well	inhibitio n rate	n n	Accumulation	inhibition
906	1417	12156	1730	9.091655	0.22086		0.104175	0.306157	0.895825	0.25471027
300	1339	12288	1625	9.141776	9.21240		0.118062	0.2019827	1,777764	9
100	1436	18834	1419	0.079349	0	0.09024 5	0.056531	0.083921	2.721232	8
33.3333	1572	16034	1634	0	0	0	0	0.0273897	3,721232	2
11.1111	1750	17652	1432	0	0	0.08216	0.02739	0.02739	4.693843	0,00880137
Control	Cell	15602		Blank			Dilution	3	1050	FALSE
			Inh	Inhibition effect to HasAg	ect to HB	sAg				
	Firs	Secon	H.	Inh	Inhibition rate	rte	Average	A te luminos	4	Accumulate
n(x10/10/cdl)	1	4 to 11	, Tell	First	Second	Third	ionibitio n rate		Acquantation	inhibition
906	1208	11692	1223		0.01275	, o	0.00425	0.025163	6.99575	3.02464711
300	1284	11484	1235	٥	0.03031		0.010104	0.0209125	1.985646	3.01042207
100	1289	14696	1508	٥	0	ō	o	0.010808	2.985646	0.00360695
33.3333	1503	12928	1302	0	۰	0	0	0.0108081	3.985646	6
11.1111	1179	11984	1150	0.004137	۰	9.02828	0.010808	0.010808	4.934837	0.00216783 8
Control	2118	11843		Blank	o		Dilution	en	1050	FALSE

Second betch: (Infergen)

Conceasing First Secon Philadelical Tababilician Tabab						Inhit	oftion effe	Inhibition effect to HBeAg			
Color Colo	Content cat lo	Firs	Becon	Thir	di.	lbition ra	ıţ.	Average		4	Accumulate
1278 5776 6409 0, 2000511 0, 107746 0, 128457 0, 168	n(×10'II/n1)	r well	d well	d well	First	Second We11	Third	inhibitio n rate	n n	Accumulatio n	inhibition
100 100	906	6278	6375	6408	0.200051	0.18756	0.18348	0.190367	0.274635	0.809633	0.25329050
1.0 1.0	300	7692	9092	6394	7.019877	۰	0.18527	0.066383	0.0842678	1.74125	5.04616100
1962 1964 1969 0	100	8960	7474	01190		0.04765 s	0	0.015885	0.015885	2,725365	0.00579485 6
1764 1768 1768 0	33.3333	8830	8144	9682	Q	0		0	0	3,725365	0
Col. 7048 104 10	11,111,11	7848	7848	7848	0	0		0	0	4.725365	0
First Secondary This bill before to Range Paris Secondary This Secondary	Control	Ce11	7848		Blank	o		Dilution	8	IC50	FMLSE
Fig. Secon This Initializin care Second Fig. Initializin care Second Fig. Second Third Initializin care Second Third Initializin care Third Initializin care Third Third				Ę	ibition eff	ect to MBs	Ang				
v d d v rest second rest n n n rest rest	Concess on to	Fixe	Secon	Thir	I ah	thition ra	ę.	Average		1-	Accumulate
1256 1256 1277 0.03871 0.07385 0.04318 0.041004 0.15287 1.041004 0.15287 0.00318 0.15287 0.00318 0.15287 0	n(x[0,10/m])	, seli	well well	, į	First well	Sacond well	Third	inhibitio n zate	n.	Accumulation	inhibition rate
115 1210 1277 0.195097 0.00335 0 0.13267 1277 0.152697 0.00325 0 0.003267 1277 0.152697 0.1	006	1236	12268	1227		0.04365 5	0.04318	0.041004	0.140162	0.958996	0.12751773
224 13468 1368 1368 0.023623 0 0.003614	300	1159	12708	1371		0.00935	0	0.035267	0.0991581	1,923709	0.0490186
1281 11346 1244 0.016526 0.11552 0.02593 0.053996 1282 1282 0.053996 0.053996	100	1244	13468	1398	0.029623		0	0.009874	0.063871	2.913834	0.02144964
1282 12828 1282 0 0 0 0 0	33.33333	1261	11346	1244	0.016526	9.11552	0.02993	0.053996	0.0539965	3.859838	0.01379630
	11.1111	1282	12826	1282		0	o		0	4.859838	Q
Control Cell 12828 Blank 0 Bilution 3	Control	Cell	12828		Blank			Bilution	3	1C50	FALSE

Third batch: (Infergen[®])

						-				
					Jrhit	ition eff	Inhibition effect to abong	51		
	1	1		Inhi	Inhibition rate	ate	Average	Parisment ar i		Accommutat
O'TU/el)	1111	well	112	Pirst well	Second	Third	inhibitio n rate	e	Accumulati on	inhibitio n rate
906	7240	6642	62.5	0.06459	0.1418	0.20439	0.136951	0.217399	0.853049	35
300	11072	8786	2069.		0	0.10826	0.03609	0.0804479	1.82696	64
100	7016	9726	7552	0.09354	D	9.02428	0.039276	0.044358	2,787683	17
33.3333	7622	9988	8676	0.01524 5			0.005082	0.0050818	3,782601	7.1
11.1111	7740	7740	7740	0	o		۰	0	4.782601	0
Control	Cell	7740		Blank			Dilution	3	ICSO	FALSE
		Ŧ	phibition	Inhibition effect to HBsAq	o HBsAq			İ		
	1		1	Inh	Inhibition rate	ate	Average	Acres ar.		Accumulat
O'IU/al)	110	Second well	well	First	Second	Third	inhibitio n rate	6	Accumulati on	inhibitio n rate
906	11048	11856	11902	0.04775		٥	0.015917	0.015917	0.984083	0.0159167 96
300	13454	12896	11798	۰		٥	0	0	1.984083	0
100	12846	13160	32546					0	2.984083	0
33.3333	12680	12458	12360	0			0	0	3.984083	٥
11.11111	11602	11602	11602	o	0		0	0	4.984083	0
Control	Cell	11602		Blank			Dilution	3	IC50	FALSE

ilbehg: Average IC50: 0 SD: 0

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EXAMPLE 5

Preparation of rSIFN-co

Preparation of lyophilized injection

Lyophilized powder

Stock Solution of 34.5 µg/ml

rSIFN-co

in freeze dryer.

10mmol/L

PB (pH7.0) Glycine

0.4mol/L

Preparation technique: Weigh materials according to recipe. Dissolve with sterile and pyrogen-free water. Filter through 0.22 mm membrane to de-bacterialize, preserve at 6-10°C. Fill in vials after affirming it is sterile and pyrogen-free, 0.3 ml /vial or 0.5 ml/vial, and lyophilize

Preparation of liquid injection

Solution

Stock Solution of 34.5 µg/ml

rSIFN-co

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20

25mmol/L PB (pH7.0)

NaCl

0.lmol/L

Preparation: Weigh materials according to recipe. Add to desired level with sterile and pyrogen-free water. Filter through 0.22µm membrane to de-bacterialize, preserve at 6-10°C. Fill in airtight vial after affirming it is sterile and non-pyrogen at 0.3 ml /vial or 0.5 ml/vial. Storage at 2-10°C, and protect from light.

EXAMPLE 6

Armte Toxicity of rSIFN-co

Treat mice with large dose (150µg/kg, equal to 1000 times 25 of the normal dose per kilc used in treatment of adult patients) of rSTFN-co at one time by intramuscular injection. Then, observe and record their deaths and toxic reactions. Results show that: 24 hours efter injection, no abnormal reaction had been recorded. The organs of the animals which had been selected to be killed also had no signs of abnormal changes. Those remaining mice were all kept alive and were normal after two weeks. The weights of mice in the experimental group and control group all increased, and the ratio of increase had no obvious difference between the two groups (F>0.05) according to their weights on the fourteenth day. No abnormal changes were seen from the main organs of those mice after two weeks.

1. Experimental material

1.1 Animals

40 healthy adult mice, weighing 18-22g, helf male and half female, qualified by Sichuan experiment animal control center.

2.2 Medicines

rSIFN-co (Provided by Sichuan Muiyang Life-engineering 0 Ltd.) sterilized solution, 0.15 mg/ml, Lot: 981201 rSIFN-co was administered i.m. in saline.

2. Method

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Separate the 40 mice into two groups randomly, one for experimental medicine, another for control. Inject medicines or saline at the same ratio (0.1 ml/10 g) through muscle to each mouse according to which group they belong. (150 µg/kg of rSIFN-co for experimental group; and saline for control group). After injection, observe and record acute toxicity shown in mice. Kill half of the mice (male and female each half) to check whether there were any abnormal pathologic changes in their main organs, such as heart, spleen, liver, lung, kidney, adrenal gland, stomach, duodenum, etc. after 24 hours. Those remains were kept and observed until the fourteenth day. Weigh all mice, kill them, and then observe the appearance of the organs listed above to see if there are any abnormalities. Take

pathological tissue and examine it, using the examination to assess the difference in weight increases in the two groups.

Results 5 Results show that there was no acute toxicity seen after all mice were treated with i.m. rSIFN-co with 150 µg/kg at a time, equal to 1000 times the normal dose per kilo used in treatment of adult patients. In the 14 days after injection, all mice lived well. They ste, drank, exercised, and excretad normally and showed normal hair conditions. None of them died. The observation of the main organs of the randomly selected mice shows no abnormal changes 24 hours after injection. 14 days after injection, all remaining mice were killed. Autopsies also showed no 15 changes. The weights of mice in the two groups all increased, but no obvious difference was shown when accessed with statistic method (p > 0.05). See Table 1:

Table 1 Influence to weights of mice after injection of

rSIFN-co					
Group	Dose	Animal	Weights before injection (g)	Weights after injection (g)	Increased value of weights (g)
Control	0	20	19.8 ±	30.8 ±	11.0 ± 2.9
rSIFN-co	150	20	19.4 ±	32.1 ±	12.7 ±

3. Conclusion

Under conditions of this experiment, there were no toxic reactions in all mice after injection of rSIFN-oo with 150 mg/kg. The conclusion can be reached that the maximum tolerable dose of i.m. in mice is 150 mg/kg, which is equal to 1000 times the normal dose per kilo used in treatment of adult patients.

EXAMPLE 7

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The clinic effects of recombinant super-compound interferon (rSIFM-co)

The recombinant super-compound interferon (rSIFN-co) is an invention for viral disease thorapy, especially for hepatitis. Meanwhile, it can inhibit the activity of EB viruses, VSV, Herpes simplex viruses, connaviruses, measles viruses et al. Using Wish cells /VSV system as the assay for anti-virus activity, the results showed that: the other rIFN, was 9×10° 10/mg, Intron A was 2.0×10° Id/mg and rSIFN-co was 9×10° ID/mg. The anti-viral activity of rSIFN-co is much higher than those of the former two.

Under the permission of the State Food and Drug Administration (SFDA), People's Republic of China, the clinical trials have taken place in Western China Mospital of Sichman University, the Second Respital of Chongqing Medical University and the First Affiliated Hospital of Zhejiang University School of Medicine since February 2003. The clinical treatment which focuses on the Hepstitis B is conducted under the guidance of the mutilcanter, double-blind random test. IFN-olb was used as control, and the primary results showed the following:

The affect of xSIFN-co compared with IFN- α lb in the treatment of chronic active Hepatitis B

- Standard of patients selection: The standard 1-4 are effective to both treatment with rSIFN-co (9yg) and IFN-db (SMU, 50yg), and the standard 1-5 are for rSIFN-co (15yg) treatment.
 - 1). Age: 18-65
- HBBAG test positive last over six months, HBeAg test positive, PCR assey, HBV-DNA copies ≥10³/ml
 - 3). ALT ≥ two times of the normal value
 - 4). Never received IFN treatment; or those received the Lamividine treatment but failed or relapsed
- 5) Once received other IFNs (3MU or 5MU) treatment six months ago, following the standard of SFDA but failed or relapsed

2. Evaluation of the effects:

In reference to the recommendations from the Tenth China National Committee of Virus Hepatitis and Hepatopathy, the effects were divided into three degrees according to the ALT level, HBV-DNA and HBeAy tests.

Response: ALT normal level, HBV-DNA negative, HBeAg

Partial response: ALT normal level, HEV-DNA or HBeAg

Non response: ALT, HBV-DNA and HBeAg unchanged

The response and partial response groups consider as

effective cases.

3. Results of clinic trial:

Group A: treatment with rSIFN-co(9µg)

Group B: treatment with IFN-clb (5MU, 50 µg) HBV-DNA Heptal KBsAg **BBeAg** Transfe Transfe Transfe functio Effect1 cas r to Per gro Medicine r to r to negati⊽ iod ้นอ ėŝ Recover Rate negativ negativ rate e rate e rate rate 28.12 37.50 84.38 rSIFN-46.88 9.38 32 (27) (12) (15) (3) (9) co (9µg) 12 IFN-alb wee 9.38 15.62 56.25 21.88 0.00 32 (5MU, 50 k (0) (3) (5) (18) (7) pg) 34.38 90.62 7.81 25.00 rSIFN-54.69 64 16-(58) (22) co (9µg) (35)(5) (16) 24 IFN-alb wee 18.75 78.13 25.00 0.00 9.38 (5MO, 50 (12) (50) (16)(0) (6) μq)

In Group C, the cases were chronic active Mepatitis B treatment with other IFNs (3MD or 5MD) before but failed or relapsed and treated with rSIRM-co (15 µg), subcutaneous injection, every one day, last 24 weeks. The total cases are 13. After 12 weeks treatment, 7 of 13 (53.85%) were effective. 3 of 13 (23.08%) HBeAg transferred to negative: 7 of 13(53.85%) HBV-DRA transferred to negative: 11 of 13

(84.62%) hepal functions recovered to normal.

- 4. The side effects of rSIFN-co compared with IFN-olb in the treatment
- 5 The side effects of JFN include fever, nausea, myalgia, anorexia, hair lose, leucopenia and thrombocytopenia, etc. The meximum dose of IFN-alb is 5MIU per time; the routine dose is 3 MIU. When taken the routine dose, 90% patients have I- II degree (WHO standard) side effects. They are 10 fever lower than 38°C, nausea, myalgia, anorexia, etc. When taken at maximum dose, the rate of side effects do not rise obviously, but are more serious. The maximum dose of rSIFMco is 24µg, subcutaneous injection, every one day for 3 months. The routine dose is .9µg. When routine doses were 15 used, less than 50% patients have I-II degree (WHO standard) side effects, including fever below 38°C, nausea, myalgia, anorexia, leucopenia and thrombocytopenia slightly. With maximum dosage, about 50% patients suffered from leucopenia and thrombocytopenia after using rSIFM-co one month, but 20 those side effects would disappear after stopping treatment

for one week. It is safe for continue use. The observations of rSIFM-co treat Hepatitis C

- 1. Standard of patient's selection
- 5 1) age: 18-65
 - 2) HCV antibody positive
 - 3) ALT \geq 1.5 times of the normal value, last more than 6 months
 - 2. Evaluation of the offects:
- 0 Referring to the standard of Infergen® for treatment of Hepatitis C and according to the ALT lavel and HCV-RNA test, divided the effects into three degree: Response: ALT normal level, HCV-RNA negative Partial response: ALT normal level, HCV-RNA unchanged
- 35 Non response: ALT and HCV-RNA unchanged

3. Effects in clinic

The clinical trial was done at the same time with Hepatitis B treatment. 46 cases received the treatment, 9 µg each time, subcutaneous injection, every day for 24 weeks. After treatment, 26 of 46 (56.52%) have obvious effects, 12 of 46 (66.08%) HCV-RNA transferred to negative, 26 of 46 (56.52%) hepal functions recovered to normal.

When used in this specification and claims, the terms
"comprises" and "comprising" and variations thereof mean
that the specified features, steps or integers are
included. The terms are not to be interpreted to exclude
the presence of other features, steps or components.

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What is claimed is:

- A recombinant super-compound interferon or a functional equivalent thereof with changed spatial configuration and improved efficacy.
- 2. The interferon of claim 1, wherein the interferon is either α , β , or ω .
- The interferon of claim 1, wherein the interferon has higher efficacy than the interferon described in U.S. Patent Nos. 4,695,623 or 4,897,471, or with unique secondary or tertiary structure.
- 15 4. The super-compound interferon of claim 1, wherein the spatial configuration change is the result of changes of its production process.
 - A super-compound interferon of claim 1, produced by a high efficiency expression system which uses a special promoter, or the promoter is Pas.
 - The super-compound interferon of claim 4, wherein its gene is artificially synthesized cDNA with adjustment of its sequence from the wild-type according to codon preference of E. coli.
 - The super-compound interferon of claim 1, which possesses anti-viral or anti-tumor activity.
 - 8. The super-compound interferon of claim 7, wherein the virus diseases comprises Hepatitis A, Hepatitis B, Hepatitis C, other types of hepatitis, infections caused by Epstein-Barr virus. Cytomegalovirus, herpes simplex viruses, other herpes viruses, pepovaviruses, poxviruses, picornaviruses, adenoviruses, rihnoviruses, human T cell leuksemia viruses I, human

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T cell leukaemia viruses II, or human T cell leukemia viruses III.

- The super-compound interferon of claim 8, which directly inhibits the DNA duplication and secretion of HBsAg and HBeAg of Hepatitis B Virus.
- 10. An artificial gene codes for the super-compound interferon or its equivalent of claim 1.
 - 11. A vector comprising the gene of claim 10, an expression system comprising the said vector, or a host cell comprising the said vector.
- 12. A process for production of recombinant supercompound interferon comprising introducing an artificial gene with selected codon preference into an appropriate boxt, culturing said introduced host in an appropriate condition for the expression of said compound interferon and harvesting the expressed compound interferon.
- 13. The process for production of claim 12, comprising of super-compound interferon 25 extraction fermentation broth, collection of inclusion body, denaturation and renaturation of the harvested protein, or wherein the process maintains the high efficacy even when the super-compound interferon is used with an agent and in a particular concentration, 30 or comprising separation and purification of the super-compound interferon, or comprising lyophilization of the purified super-compound interferon, or comprising production of liquid injection of super-compound interferon. 35
 - 14. The produced super-compound interferon by the process

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of any of the claims 12-13.

- 15. A composition comprising the recombinant supercompound interferon of claim 1 and a suitable carrier.
- 16. A pharmaceutical composition comprising recombinant super-compound interferon of claim 1 and a pharmaceutically acceptable carrier.
- 17. A method for treating viral diseases or tumor in a subject comprising administering to the subject an effective amount of the super-compound interferon of claim 1.
- 18. The method of claim 17 wherein the viral diseases is 15 Repatitis A, Repatitis B, Repatitis C, other types of hepatitis, infections of viruses caused by Epstein-Barr virus, Cytomegalovirus, herpes simplex viruses, or other type of herpes viruses, papovaviruses, 20 poxviruses. picornaviruses. rihnoviruses, human T cell leukaemia viruses I, or human T cell leukaemia viruses II, or human T cell
- 25 19. The method of claim 17 wherein super-compound interferon was administered via oral, vein injection, muscle injection, peritoneal injection, subcutaneous injection, nasal, mucosal administration, by 30 inhalation via an inspirator.

leukemia virus III.

20. The method of claim 17 wherein super-compound interferon was administered following the protocol of injection 9 mg or 15 mg per day, 3 times a week, total 24 weeks.

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Figure 1

S' 11 21 31 41 51
+1 N C D L P O T H S L E N R R A L L L A
1 ATTERCORLOS TOCOGRAMOS COMPOSOR GETANGGETH STEETITIBAT COTRECTMENT
TANDICIDITES ACROSOTOTIS GETCASCANC CONTINUOUS COMPANANTA CRUCIANOCON

5° N. R. I. S. P. F. S. D. L. X. D. M. P. F. F. F. F. CARTACTOR STATEMENT CONTINUOUS GENERAL CHICAGOTTI CONTINUOUS GENERAL CHICAGOTTI CONTINUOUS CHICAGOTT

5' 133 141 151 161 171
10 E E F D G N G F O K N D A 1 S Y L H E
121 CASSAMANT TOSAGOSTAN CONSTITUOUS AMESTOAGE CHATCOOCT TOTOCNOSIAN
STOCTICITA ANCTEGOCHT GETCANGSTC TITTCHASTOC GATAGASOON AGAGSTSCHT

S' B 1 O T F Y L F S T M 21 231 251 51 B 1 O T F Y L F S T M 221 A 31 O T F Y L F S T M A W 0 E 181 ATRACOACO AGACTICA COTATICOS ACCAMANOS TOCIOGOSTOS TISSOMOSTICA CONTROL TRACOACIONA CONTROL TRACOACIONA CONTROL TRACOACIONA ACCORDINATION CONTROL TRACOACIONA ACCORDINATION CONTROL TRACOACIONA ACCORDINATION CONTROL TRACOACIONA ACCORDINATION CONTROL TRACOACIONA CONTRO

5' 251 261 271 281 271 281 271 41 5 L E A C 17 F L Y O O L N D L E A C 241 TOCHOTEGRA AMANITORI ACCOMANTE RACHMORACE TEANGRACH GAMAGET
5' 311 321 331 341 351
11 0 EV 0 7 E T P L N N 0 S 1 L A
301 STATICAGETOS TISTAGORIAS ACTIOTITOS GEDACTACT YEDAMORIAS GETASORIOS TISTAGORIAS ACTIOTITOS GEDACTACT YEDAMORIAS GENASORIOSA

5' 371 S81 391 401 411
+1 Y K K Y F Q R I T L Y L T E K K Y S P C
381 STEAMANT ACTROACES TATCHESSAG ATSCACTOR AMMANTA (TOCCOTTOE
CONSTITUTO A SAGGIOSE ATSCACTOR THITTITAL FASSSACKOS

5' 491 501 +1 E R L R R K E # 481 EAROGICIGO BIOGRAPAGA ATAA CTIGGAGAGO CAGGATITOT TATE

Figure 2

5° 11 21 31 41 51

+1 M C D L P Q T H S L G N R R A L I L A

1 ATISTICATI TACKTOWN TONTICOTH GRANDIST GESCHOT TOTICSTEED
TROPOLICIA ATISTICTI SATAVONA CONTINUONS GESPECTA ANGENTS

F 71 81 91 1 11
+1 Q M R R I S P F S C L K D R H D F G F P
81 OPERIOR CHARTOCT CHARACTIC GOVERNOUS GOVERNMENT CONTINUES CHARACTIC CHARAC

5° 31 41 51 61 71 +I O E E F D G N Q F Q K A Q A I S V L H E 121 CAMPANDAT TOORTEDDA COMPTONE MACTICAGO CONTOCTES ACTICAGON COTOTOTO ACCIDICATO CONTOCTO OTRAGENO TOMOSPICAT

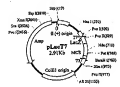
5 91 1 11 21 31 +1 M I Q Q T F N L F S T K D S S A A W D E 181. ADRICOMA CARGOTTON CITETTOC ADTAMASON GETCHORGE THRESPAGNA TACAGRIETE TORGAMENT GENOMAGE PRITITIONS CONCINCENT

5' 51 51 71 81 91

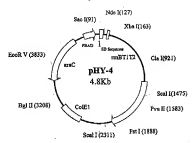
+) S L L E K F Y T E L Y Q Q L N D L E A C
241 ACCITICISTA AGNATICIA OPCIGNACIS TRANSCACIS GRANDATICI
TICIANOSIC TICITONAST GENETISIC ATACISTICS ACTICISTACIS

5" 11 21 31 41 51 +1 V I Q E V G V E E T P L M N V D S I L A 301 GRANDOGA MAGNIGGIA MAGNIGAT CERDIGATIA ACCIGACIO TATTURIGISA OUTRAGISCO TIDMODOA TOTTORIORA GEOGRAFIA MAGNICAT

Figure 3







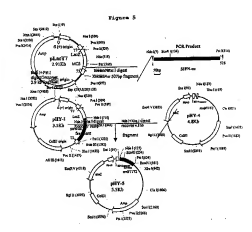
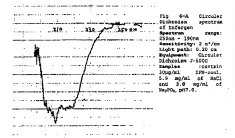


Figure 6-A

Circular Dichroism spectra Tested by Analysis and Measument Center of Sichuan University.





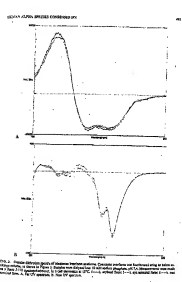


Fig 6-B Circular Dichroism spectrum of Infergen From Reference[Journal of Interferon and cytokine Research. 16:489-499(1996)]

Figure 6-C



Fig 6-C Circular Dichroism spectrum of fSFN-co Spectrum yrange 320mm-250mm Sensitivity 2 m*/com Light path 2cm Zutiperate Circular Dichroism 1-500C Samples: comain 0.5mg/ml (SFN-co, 5.9 mg/ml of NaCl and 3.8 mg/ml of NacPOs, pH7.0.

Figure 6-D

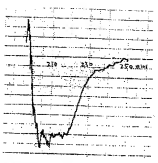


Fig 6-D Circular Dichroism spectrum of rSIFN-co Spectrum range; 250nm - 190nm Sensitivity: 2 m°/cm Light path: 0.20 cm

Equipment: Circular Dichroism J-500C Samples :contain 30µg/ml rSiFN-co, 5.9 mg/ml of NaCl and 3.8 mg/ml of Na₂PO₄ pH7.0.